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NEWS

TERMINAL (ENTER 1, 2, 3, OR ?):2

\* \* \* \* \* \* \* \* \* \* Welcome to STN International

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NEWS 5 May 12 EXTEND option available in structure searching NEWS 6 May 12 Polymer links for the POLYLINK command completed in REGISTRY

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NEWS 9 May 27 CAplus super roles and document types searchable in REGISTRY

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NEWS 13 Jun 28 ANTE, AQUALINE, BIOENG, CIVILENG, ENVIROENG, MECHENG, and WATER from CSA now available on STN(R)

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NEWS WWW CAS World Wide Web Site (general information)

Enter NEWS followed by the item number or name to see news on that specific topic.

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=> file reg

COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.21 0.21

FULL ESTIMATED COST

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provided by InfoChem.

STRUCTURE FILE UPDATES: 28 JUN 2004 HIGHEST RN 700803-86-7 DICTIONARY FILE UPDATES: 28 JUN 2004 HIGHEST RN 700803-86-7

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2004

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

```
=> e isophorone diisocyanate/cn
                   ISOPHORONE DIAMINE-VESTICOAT UT 647 COPOLYMER/CN
             1
                   ISOPHORONE DIICYANATE-UPICACOAT GV 150 COPOLYMER/CN
E2
             1
E3
             1 --> ISOPHORONE DIISOCYANATE/CN
                   ISOPHORONE DIISOCYANATE 2-HYDROXYPROPYL ACRYLATE (1:2) ADDUC
E4
E5
             1
                   ISOPHORONE DIISOCYANATE ADDUCT WITH 2-ETHYLHEXANOL AND N,N-D
                   IMETHYLAMINOETHANOL/CN
E.6
             1
                   ISOPHORONE DIISOCYANATE ADDUCT WITH TRIETHYLENE GLYCOL MONOM
                   ETHYL ETHER AND N, N-DIMETHYLAMINOETHANOL/CN
                   ISOPHORONE DIISOCYANATE CAPROLACTAM ADDUCT (1:2)/CN
E7
             7
                   ISOPHORONE DIISOCYANATE CYCLIC TRIMER/CN
             1
E8
                   ISOPHORONE DIISOCYANATE DIUREA WITH OCTADECYLAMINE/CN
             1
E9
                   ISOPHORONE DIISOCYANATE DIURETHANE WITH 4-OCTYLPHENOL ETHOXY
             1
E10
                   LATE/CN
E11
             1
                   ISOPHORONE DIISOCYANATE DIURETHANE WITH OCTADECYL ALCOHOL/CN
                   ISOPHORONE DIISOCYANATE DIURETHANE WITH TETRAHYDROABIETYL AL
E12
             1
                   COHOL/CN
=> e3
             1 "ISOPHORONE DIISOCYANATE"/CN
L1
=> d 11
    ANSWER 1 OF 1 REGISTRY COPYRIGHT 2004 ACS on STN
T.1
     4098-71-9 REGISTRY
RN
     Cyclohexane, 5-isocyanato-1-(isocyanatomethyl)-1,3,3-trimethyl- (9CI) (CA
CN
     INDEX NAME)
OTHER CA INDEX NAMES:
     Isocyanic acid, methylene(3,5,5-trimethyl-3,1-cyclohexylene) ester (7CI,
     8CI)
OTHER NAMES:
CN
     1,3,3-Trimethyl-1-(isocyanatomethyl)-5-isocyanatocyclohexane
     1-(Isocyanatomethyl)-5-isocyanato-1,3,3-trimethylcyclohexane
CN
     1-Isocyanato-3,3,5-trimethyl-5-(isocyanatomethyl)cyclohexane
CN
     1-Isocyanato-3-(isocyanatomethyl)-3,5,5-trimethylcyclohexane
CN
CN
     1-Isocyanato-5-(isocyanatomethyl)-3,3,5-trimethylcyclohexane
CN
     3,3,5-Trimethyl-5-(isocyanatomethyl)cyclohexyl isocyanate
     3-(Isocyanatomethyl)-3,5,5-trimethylcyclohexyl isocyanate
CN
     5-Isocyanato-1-(isocyanatomethyl)-1,3,3-trimethylcyclohexane
CN
CN
     IPDI
CN
     Isophorone diisocyanate
FS
     3D CONCORD
     124961-52-0, 63793-40-8, 129212-17-5, 101701-80-8, 102771-74-4,
DR
```

105439-02-9, 66708-07-4, 50974-99-7, 74091-63-7, 74520-92-6, 70936-97-9,

146282-59-9, 146665-38-5, 149579-36-2, 88778-74-9, 26602-93-7, 52985-93-0, 110648-35-6, 111093-75-5, 194936-84-0

MF C12 H18 N2 O2

CI COM

LC STN Files: AGRICOLA, ANABSTR, BEILSTEIN\*, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS, CHEMINFORMRX, CHEMLIST, CHEMSAFE, CIN, CSCHEM, CSNB, DIOGENES, DIPPR\*, EMBASE, HSDB\*, IFICDB, IFIPAT, IFIUDB, MEDLINE, MSDS-OHS, NIOSHTIC, PIRA, PROMT, RTECS\*, SPECINFO, TOXCENTER, ULIDAT, USPAT2, USPATFULL (\*File contains numerically searchable property data)

Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

DT.CA CAplus document type: Conference; Journal; Patent; Report

RL.P Roles from patents: ANST (Analytical study); BIOL (Biological study);
MSC (Miscellaneous); OCCU (Occurrence); PREP (Preparation); PROC
(Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses);
NORL (No role in record)

RLD.P Roles for non-specific derivatives from patents: ANST (Analytical study); BIOL (Biological study); FORM (Formation, nonpreparative); MSC (Miscellaneous); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)

RL.NP Roles from non-patents: ANST (Analytical study); BIOL (Biological study); MSC (Miscellaneous); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)

RLD.NP Roles for non-specific derivatives from non-patents: ANST (Analytical study); BIOL (Biological study); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

4334 REFERENCES IN FILE CA (1907 TO DATE)
3235 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
4340 REFERENCES IN FILE CAPLUS (1907 TO DATE)
2 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

=> file reg COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 8.30 8.51

FULL ESTIMATED COST

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STRUCTURE FILE UPDATES: 28 JUN 2004 HIGHEST RN 700803-86-7

DICTIONARY FILE UPDATES: 28 JUN 2004 HIGHEST RN 700803-86-7

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2004

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

=> bromine

L2 2357 BROMINE

=> isocyanate

17997 ISOCYANATE 5 ISOCYANATES

L3 17997 ISOCYANATE

(ISOCYANATE OR ISOCYANATES)

=> 12 and 13

L4 7 L2 AND L3

=> d 14 1-7 ti

'TI' IS NOT A VALID FORMAT FOR FILE 'REGISTRY'

The following are valid formats:

Substance information can be displayed by requesting individual fields or predefined formats. The predefined substance formats are: (RN = CAS Registry Number)

REG - RN

SAM - Index Name, MF, and structure - no RN FIDE - All substance data, except sequence data

IDE - FIDE, but only 50 names
SQIDE - IDE, plus sequence data

SQIDE3 - Same as SQIDE, but 3-letter amino acid codes are used

SQD - Protein sequence data, includes RN

SQD3 - Same as SQD, but 3-letter amino acid codes are used SQN - Protein sequence name information, includes RN

CALC - Table of calculated properties

EPROP - Table of experimental properties

PROP - EPROP and CALC

Any CA File format may be combined with any substance format to obtain CA references citing the substance. The substance formats must be cited first. The CA File predefined formats are:

ABS -- Abstract

APPS -- Application and Priority Information

BIB -- CA Accession Number, plus Bibliographic Data

CAN -- CA Accession Number

CBIB -- CA Accession Number, plus Bibliographic Data (compressed)

IND -- Index Data

IPC -- International Patent Classification

PATS -- PI, SO

STD -- BIB, IPC, and NCL

```
IABS -- ABS, indented, with text labels IBIB -- BIB, indented, with text labels
```

ISTD -- STD format, indented

OBIB ----- AN, plus Bibliographic Data (original)

OIBIB ----- OBIB, indented with text labels

SBIB ----- BIB, no citations SIBIB ----- IBIB, no citations

The ALL format gives FIDE BIB ABS IND RE, plus sequence data when it is available.

The MAX format is the same as ALL.

The IALL format is the same as ALL with BIB ABS and IND indented, with text labels.

For additional information, please consult the following help messages:

HELP DFIELDS -- To see a complete list of individual display fields.
HELP FORMATS -- To see detailed descriptions of the predefined formats.
ENTER DISPLAY FORMAT (IDE):file reg
'FILE' IS NOT A VALID FORMAT FOR FILE 'REGISTRY'

The following are valid formats:

Substance information can be displayed by requesting individual fields or predefined formats. The predefined substance formats are: (RN = CAS Registry Number)

REG - RN

SAM - Index Name, MF, and structure - no RN FIDE - All substance data, except sequence data

IDE - FIDE, but only 50 names
SQIDE - IDE, plus sequence data

SQIDE3 - Same as SQIDE, but 3-letter amino acid codes are used

SQD - Protein sequence data, includes RN

SQD3 - Same as SQD, but 3-letter amino acid codes are used SQN - Protein sequence name information, includes RN

CALC - Table of calculated properties EPROP - Table of experimental properties

PROP - EPROP and CALC

Any CA File format may be combined with any substance format to obtain CA references citing the substance. The substance formats must be cited first. The CA File predefined formats are:

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APPS -- Application and Priority Information

BIB -- CA Accession Number, plus Bibliographic Data

CAN -- CA Accession Number

CBIB -- CA Accession Number, plus Bibliographic Data (compressed)

IND -- Index Data

IPC -- International Patent Classification

PATS -- PI, SO

STD -- BIB, IPC, and NCL

IABS -- ABS, indented, with text labels

IBIB -- BIB, indented, with text labels

ISTD -- STD format, indented

OBIB ----- AN, plus Bibliographic Data (original) OIBIB ---- OBIB, indented with text labels

SBIB ----- BIB, no citations SIBIB ----- IBIB, no citations

The ALL format gives FIDE BIB ABS IND RE, plus sequence data when it is available.

The MAX format is the same as ALL.

The IALL format is the same as ALL with BIB ABS and IND indented. with text labels.

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HELP DFIELDS -- To see a complete list of individual display fields. HELP FORMATS -- To see detailed descriptions of the predefined formats. ENTER DISPLAY FORMAT (IDE):end

=> file caplus

COST IN U.S. DOLLARS

SINCE FILE ENTRY

TOTAL SESSION 18.21

FULL ESTIMATED COST

9.70

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FILE COVERS 1907 - 29 Jun 2004 VOL 141 ISS 1 FILE LAST UPDATED: 28 Jun 2004 (20040628/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> bromine

47316 BROMINE

152 BROMINES

L5 47408 BROMINE

(BROMINE OR BROMINES)

=> isocyanate

58621 ISOCYANATE

20235 ISOCYANATES

66442 ISOCYANATE L6

(ISOCYANATE OR ISOCYANATES)

=> 15 and 16

197 L5 AND L6

=> 15(1)16

L8 67 L5(L)L6

=> phosgene

12345 PHOSGENE

35 PHOSGENES

L9 12353 PHOSGENE

(PHOSGENE OR PHOSGENES)

=> 17 and 19

L10 11 L7 AND L9

=> d 119 1-11 ti

L19 NOT FOUND

The L-number entered has not been defined in this session, or it has been deleted. To see the L-numbers currently defined in this session, enter DISPLAY HISTORY at an arrow prompt (=>).

=> d 110 1-11 ti

- L10 ANSWER 1 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Integrated process for preparing **phosgene** and from it **isocyanates**
- L10 ANSWER 2 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Threshold quantity criteria for risk management programs: recommendations for toxic releases
- L10 ANSWER 3 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI The development of acute exposure guideline levels for hazardous substances
- L10 ANSWER 4 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Accidental release prevention
- L10 ANSWER 5 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI The Use of Benchmark Dose Methodology with Acute Inhalation Lethality Data
- L10 ANSWER 6 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Incorporation of potential for multi-media exposure into chemical hazard scores for pollution prevention
- L10 ANSWER 7 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Management & design of process exhaust systems in an I/C manufacturing environment for emission minimization
- L10 ANSWER 8 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Performance-oriented packaging standards; changes to classification, hazard communication, packaging and handling requirements based on UN standards and agency initiative
- L10 ANSWER 9 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Air contaminants
- L10 ANSWER 10 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of aliphatic diisocyanates without using phosgene
- L10 ANSWER 11 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI dl-7-Azidocephalosporins

```
ANSWER 1 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
     Integrated process for preparing phosque and from it
     isocyanates
AN
     2004:100871 CAPLUS
DN
     140:165773
TI
     Integrated process for preparing phosgene and from it
     Walsdorff, Christian; Fiene, Martin; Strofer, Eckhard; Harth, Klaus;
IN
     Jacobs, Jan D.; Deberdt, Filip
PA
     BASF Aktiengesellschaft, Germany
     U.S. Pat. Appl. Publ., 9 pp.
     CODEN: USXXCO
DT
     Patent
LΑ
    English
FAN.CNT 1
     PATENT NO.
                     KIND DATE
                                         APPLICATION NO. DATE
     _____
                                         -----
                                                         _____
PΙ
    US 2004024244
                           20040205
                      A1
                                         US 2002-227865
                                                          20020827
                                         DE 2002-10235476A 20020802
    DE 10235476
                      A1
                           20040212
                                         DE 2002-10235476 20020802
    WO 2004014845
                      A1
                           20040219
                                         WO 2003-EP8430 20030730
            PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN,
            TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY,
            KG, KZ, MD, RU
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG,
            CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC,
            NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ,
            GW, ML, MR, NE, SN, TD, TG
                                         DE 2002-10235476A 20020802
AΒ
    A process for preparing organic isocyanates comprises: (a) making
    available a first partial amount of chlorine, with the chlorine of this
    first partial amount having a content of free and bound bromine
    and iodine of <400 ppm; (b) making available a second partial amount of
    chlorine; (c) reacting the first and second partial amts. of chlorine with
    carbon monoxide to form phosgene; (d) reacting the
    phosgene from step (c) with one or more primary amines to form the
    corresponding isocyanates and hydrogen chloride; (e) separating off
    and, if necessary, purifying the isocyanates formed in step (d);
     (f) separating off and, if necessary, purifying the hydrogen chloride formed in
    step (d); (g) catalytically oxidizing at least part of the hydrogen
    chloride separated off in step (e) by means of oxygen to form chlorine; (h)
    separating off the chlorine formed in step (g) and using at least a partial
    amount of the chlorine which has been separated off as second partial amount of
    chlorine in step (b). A process flow diagram is presented.
=> d his
     (FILE 'HOME' ENTERED AT 12:35:23 ON 29 JUN 2004)
    FILE 'REGISTRY' ENTERED AT 12:35:51 ON 29 JUN 2004
               E ISOPHORONE DIISOCYANATE/CN
L1
             1 E3
    FILE 'REGISTRY' ENTERED AT 12:38:33 ON 29 JUN 2004
L2
          2357 BROMINE
```

```
L3
          17997 ISOCYANATE
L4
              7 L2 AND L3
     FILE 'CAPLUS' ENTERED AT 12:39:45 ON 29 JUN 2004
L5
          47408 BROMINE
L6
          66442 ISOCYANATE
L7
            197 L5 AND L6
L8
            67 L5(L)L6
L9
          12353 PHOSGENE
L10
            11 L7 AND L9
=> '11
MISMATCHED QUOTE ''L1'
Quotation marks (or apostrophes) must be used in pairs,
one before and one after the expression you are setting
off or masking.
=> 11
L11
          4350 L1
=> 111/prep
FIELD CODES CANNOT BE CHANGED HERE
You may have tried to apply a field code to a term that already has a
field code. You can only add a field code to a term that has no field
code appended to it.
=> 11/prep
          4350 L1
       3164945 PREP/RL
          1655 L1/PREP
                 (L1 (L) PREP/RL)
=> 19 and 112
L13
          18 L9 AND L12
=> d 113 1-18 ti
L13 ANSWER 1 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
     Improved process for the preparation of di- and triisocyanates by
     continuous phosgenation of amines
L13 ANSWER 2 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
    Production of isocyanates in the gaseous phase
L13 ANSWER 3 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
    Improved procedure for the production of mono- and oligoisocyanates by the
    phosgenation of primary amines in the presence of catalytic amounts of
    monoisocyanates
L13 ANSWER 4 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
    Evaporative removal of aliphatic and cycloaliphatic isocyanate monomers
ΤI
     from polymeric residues
L13 ANSWER 5 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
    Purification of aliphatic isocyanates
L13 ANSWER 6 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
ΤI
    Preparation of aliphatic polyisocyanates from polyamines and
    phosgene
L13 ANSWER 7 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
    Preparation of isophorone diisocyanate
```

- L13 ANSWER 8 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of diisocyanates without using phosgene
- L13 ANSWER 9 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of isophorone diisocyanate from isophoronediamine
- L13 ANSWER 10 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Process for preparing polyurethanes for coatings
- L13 ANSWER 11 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Process for making aliphatic and cycloaliphatic polyisocyanates
- L13 ANSWER 12 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Procedure for the production of (cyclo)aliphatic diisocyanates
- L13 ANSWER 13 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Manufacture of isocyanates without phosgene
- L13 ANSWER 14 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Extraction of pure diisocyanates
- L13 ANSWER 15 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Multistep process for producing 3-isocyanatomethyl-3,5,5trimethylcyclohexylisocyanate
- L13 ANSWER 16 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Polyurethane-siloxanes
- L13 ANSWER 17 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Continuous preparation of 1-isocyanato-3-(isocyanatomethyl)-3,5,5-trimethylcyclohexane
- L13 ANSWER 18 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Isocyanates
- $\Rightarrow$  d 113 5,6,9,11,12,17,18 ti fbib abs
- L13 ANSWER 5 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Purification of aliphatic isocyanates
- AN 1996:35267 CAPLUS
- DN 124:201653
- TI Purification of aliphatic isocyanates
- IN Nozawa, Kaneo; Matsuhira, Nobuya; Naito, Taketoshi; Morinaka, Katsutoshi; Tabuchi, Toshihiko
- PA Showa Denko Kk, Japan
- SO Jpn. Kokai Tokkyo Koho, 7 pp.
- CODEN: JKXXAF
- DT Patent
- LA Japanese
- FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	JP 07278088	A2	19951024	JP 1994-74915	19940413
	JP 2915784	В2	19990705		
				JP 1994-74915	19940413

AB The process comprises heating solns. of hydrolyzable Cl-containing crude aliphatic isocyanates and inert organic solvents at 140-270°, optionally mixing the crude isocyanates with the solvents at one time or gradually, distilling away higher amts. of the solvents than the weight of the isocyanates contained for ≥2 h, and optionally distilling the isocyanates from

residue. A solution of isophorone diisocyanate (I) and 423 ppm hydrolyzable Cl in o-Cl2C6H4 was distilled at 175-185° and 560-600 mm Hg for 3 h to distill away o-Cl2C6H4, then distilled at 127° and 2 mm Hg to give 36 ppm hydrolyzable Cl-containing I with Harzen color number  $\leq$ 10.

- L13 ANSWER 6 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of aliphatic polyisocyanates from polyamines and
- AN 1995:992543 CAPLUS
- DN 124:88109
- TI Preparation of aliphatic polyisocyanates from polyamines and phosgene
- PA Mitsui Toatsu Chemicals, Inc., Japan
- SO Ger. Offen., 11 pp.

CODEN: GWXXBX

DT Patent

LA German

FAN.CNT 1

T. LTIA.	∼TA T	±						
	PAT	TENT NO.	KIND	DATE	AP	PLICATION NO.		DATE
ΡI	DE	19510259	A1	19950928	DE	1995-1951025	9	19950321
	DE	19510259	C2	19970904				
					JP	1994-50082	A	19940322
	JP	07309827	A2	19951128	JP	1995-42956		19950302
	JP	3201921	В2	20010827				
					JP	1994-50082	A	19940322
	US	5523467	A	19960604	US	1995-401807		19950310
					JΡ	1994-50082	A	19940322
	CN	1125718	А	19960703	CN	1995-104546		19950322
	CN	1062857	В	20010307				
					JP	1994-50082	A	19940322

AB In the conversion of an aliph polyamine to the polyisocyanate (e.g., m-xylylenediamine to m-xylylene diisocyanate) in an inert liquid medium, an inert gas is added to the reactor during the reaction to increase the yield of polyisocyanate and reduce the amount of phosgene required.

- L13 ANSWER 9 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of isophorone diisocyanate from isophoronediamine
- AN 1993:650204 CAPLUS
- DN 119:250204
- TI Preparation of isophorone diisocyanate from isophoronediamine
- IN Suguro, Yoshio; Kawamura, Shigenori
- PA Mitsubishi Chemical Industries Co., Ltd., Japan
- SO Jpn. Kokai Tokkyo Koho, 3 pp. CODEN: JKXXAF
- DT Patent
- LA Japanese
- FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	JP 05065265	A2	19930319		19910909 19910909

- OS CASREACT 119:250204
- AB Isophorone diisocyanate (I) is prepared by reacting isophoronediamine or its hydrochloride salt (II) with **phosgene** in an inert organic solvent followed by distilling off the solvent and treating the crude product at 160-170° in an atmospheric of inert gas. Thus, II was treated with **phosgene** in decane and the reaction mixture was distilled to give crude I, which, after gel permeation chromatog., a product containing 4.2% impurities. This product was passed through a ball filter under the introduction of nitrogen gas at 1.9 L/h at 220° for 1 h to give a

product containing 3.8% impurities.

```
L13 ANSWER 11 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
TΤ
    Process for making aliphatic and cycloaliphatic polyisocyanates
ΑN
    1990:36699 CAPLUS
DN
    112:36699
    Process for making aliphatic and cycloaliphatic polyisocyanates
TI
IN
    Thorpe, David; Smith, Richard Colin
PA
    Imperial Chemical Industries PLC, UK; ICI Americas, Inc.
    Eur. Pat. Appl., 4 pp.
SO
    CODEN: EPXXDW
DT
    Patent
    English
LΑ
FAN.CNT 2
    PATENT NO.
                   KIND DATE
                                      APPLICATION NO. DATE
    ______
                                       -----
    EP 327231
                    A1
                                      EP 1989-300602
                          19890809
PI
                                                        19890123
        R: AT, BE, CH, DE, ES, FR, GB, GR, IT, LI, NL, SE
                                       GB 1988-2674
                                                        19880205
    JP 01287128
                    A2
                        19891117
                                        JP 1989-11764
                                                        19890120
                                        GB 1988-2674
                                                        19880205
    CN 1034712
                          19890816
                                        CN 1989-100723
                     A
                                                        19890203
                                        GB 1988-2674
                                                        19880205
PATENT FAMILY INFORMATION:
FAN 1990:140004
    PATENT NO.
                    KIND DATE
                                       APPLICATION NO. DATE
                    ____
                                        _____
                          19891003
PΙ
    BR 8900512
                  Α
                                        BR 1989-512 19890203
                                        GB 1988-2674 19880205
GB 1989-1609 19890125
    (Cyclo) aliphatic polyisocyanates are prepared without COCl2 by heating diamines
AB
    with excess aromatic polyisocyanate boiling \geq 20^{\circ} above the b.p.
    of the desired isocyanate. Adding 6 g isophorone diamine over 20 min to
    400 g polymethylenepolyphenylene isocyanate (I) (62% MDI) stirred at
    120°, heating at 180° for 3 h, and distilling gave 60%
    isophorone diisocyanate and 40% I.
L13 ANSWER 12 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
TI
    Procedure for the production of (cyclo)aliphatic diisocyanates
AN
    1989:231172 CAPLUS
DN
    110:231172
ΤI
    Procedure for the production of (cyclo)aliphatic diisocyanates
    Frosch, Hans Georg; Grave, Heinrich; Stutz, Herbert; Waldau, Eckart;
IN
    Fuhrmann, Peter
    Bayer A.-G., Fed. Rep. Ger.
PΑ
    Ger. Offen., 4 pp.
SO
    CODEN: GWXXBX
DT
    Patent
LΑ
    German
FAN.CNT 1
    PATENT NO.
                    KIND DATE
                                       APPLICATION NO. DATE
                    ----
                                       -----
PΤ
    DE 3714439
                    A1
                          19881110
                                       DE 1987-3714439 19870430
    EP 289840
                    A1 19881109
                                        EP 1988-106111 19880416
    EP 289840
                    В1
                         19901017
        R: BE, DE, ES, FR, GB, IT, NL
                                        DE 1987-3714439 19870430
    US 4847408
                                        US 1988-185721 19880425
                    Α
                         19890711
                                        DE 1987-3714439 19870430
    CA 1305165
                    A1 19920714
                                        CA 1988-565025 19880425
                                        DE 1987-3714439 19870430
    JP 63280050
                    A2 19881117
                                        JP 1988-104461 19880428
```

DE 1987-3714439 19870430

OS CASREACT 110:231172; MARPAT 110:231172

AB A procedure for the preparation of OCNRNCO [R = C1-15(cyclo)aliphatic hydrocarbon

moiety] by phosgenation of the corresponding H2NRNH2 in the gas phase was characterized in that one: a) brings the gaseous diamine, optionally diluted with an inert gas or the vapors of an inert solvent, and COCl2, sep. heated to 200-600°, into reaction with each other in a cylindrical chamber at 200-600° without moving parts with the maintenance of a turbulent streaming into the reactor chamber; b) leads the gas mixture which continuously leaves the reaction chamber through an inert solvent which is kept at a temperature above the decomposition temperature of the carbamoyl chloride

corresponding to the diamine; and c) subjects the diisocyanate dissolved in the inert solvent to a distillative work-up. In this manner, COC12 and H2N(CH2)6NH2 reacted at  $400^{\circ}$  to give 98.0% OCN(CH2)6NCO.

- L13 ANSWER 17 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Continuous preparation of 1-isocyanato-3-(isocyanatomethyl)-3,5,5-trimethylcyclohexane
- AN 1975:86817 CAPLUS
- DN 82:86817
- TI Continuous preparation of 1-isocyanato-3-(isocyanatomethyl)-3,5,5-trimethylcyclohexane
- IN Schmitt, Karl; Disteldorf, Josef; Reiffer, Johannes
- PA Veba-Chemie A.-G.
- SO Ger. Offen., 10 pp. CODEN: GWXXBX
- DT Patent
- LA German
- FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	DE 2323299	A1	19741121	DE 1973-2323299	19730509
	DE 2323299	C3	19831208		
	JP 50052048	A2	19750509	JP 1974-50377	19740508
	JP 58035179	В4	19830801		
				DE 1973-2323299	19730509
	US 3916006	Α	19751028	US 1974-467976	19740508
				DE 1973-2323299	19730509

AB 1-Isocyanato-3-(isocyanatomethyl)-3,5,5-trimethylcyclohexane (I) [4098-71-9] was continuously prepared without agglomeration of the suspension by phosgenation of 1-amino-3-(aminomethyl)-3,5,5-trimethylcyclohexane (II) [2855-13-2] in an inert solvent with excess COC12 at 130-60° in previously prepared I. Thus, COC12 [75-44-5] was passed into II in PhCl containing gaseous CO2 at 30°, the mixture passed into a reactor containing I in PhCl at 130° and then into a 2nd reactor for after reaction at 130° with passing of COC12 in countercurrent through both reactors to give 98% I of Cl content 0.1%.

- L13 ANSWER 18 OF 18 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Isocyanates
- AN 1973:431672 CAPLUS
- DN 79:31672
- TI Isocyanates
- IN Edmondsen, John Neville; Hulse, Rae; Kerrigan, Vincent
- PA Imperial Chemical Industries Ltd.
- SO Ger. Offen., 17 pp.
  - CODEN: GWXXBX
- DT Patent
- LA German

FAN.CNT 1							
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE		
ΡI	DE 2249459	A1	19730419	DE 1972-2249459	19721009		
				GB 1971-47795	19711014		
	IT 967968	Α	19740311	IT 1972-29815	19720928		
				GB 1971-47795	19711014		
	NL 7213291	Α	19730417	NL 1972-13291	19721002		
				GB 1971-47795	19711014		
	BE 789809	A1	19730406	BE 1972-122871	19721006		
				GB 1971-47795	19711014		
	FR 2157485	A5	19730601	FR 1972-36382	19721013		
				GB 1971-47795	19711014		
	JP 48048419	A2	19730709	JP 1972-103161	19721014		
				GB 1971-47795	19711014		

AB Isocyanates were prepared by treating a mixture of (2) amines with COCl2. Thus, a mixture of cyclohexylamine and a polyphenylamine, prepared by reacting HCHO and PhNH2, was treated with COCl2 to give cyclohexyl isocyanate and polyphenyl polyisocyanate; a mixture of n-C8H17NH2 and n-C18H37NH2 gave n-C8H17NCO and n-C18H37NCO.

=> logoff hold COST IN U.S. DOLLARS	SINCE FILE	TOTAL SESSION				
FULL ESTIMATED COST	ENTRY 51.35	69.56				
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE ENTRY	TOTAL SESSION				
CA SUBSCRIBER PRICE -5.54 -5						

SESSION WILL BE HELD FOR 60 MINUTES
STN INTERNATIONAL SESSION SUSPENDED AT 12:50:55 ON 29 JUN 2004

Connecting via Winsock to STN

Welcome to STN International! Enter x:x

LOGINID:SSSPTA1623PAZ

## PASSWORD:

\* \* \* \* \* RECONNECTED TO STN INTERNATIONAL \* \* \* \* \* \* \* SESSION RESUMED IN FILE 'CAPLUS' AT 13:14:04 ON 29 JUN 2004 FILE 'CAPLUS' ENTERED AT 13:14:04 ON 29 JUN 2004 COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

COST IN U.S. DOLLARS	SINCE FILE	$\mathtt{TOTAL}$
	ENTRY	SESSION
FULL ESTIMATED COST	51.35	69.56
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	-5.54	-5.54

=> save temp all isocyansrch/l L# LIST L1-L13 HAS BEEN SAVED AS 'ISOCYANSRCH/L'

=> d his

```
(FILE 'HOME' ENTERED AT 12:35:23 ON 29 JUN 2004)
     FILE 'REGISTRY' ENTERED AT 12:35:51 ON 29 JUN 2004
                E ISOPHORONE DIISOCYANATE/CN
              1 E3
L1
     FILE 'REGISTRY' ENTERED AT 12:38:33 ON 29 JUN 2004
L2
           2357 BROMINE
L3
          17997 ISOCYANATE
L4
              7 L2 AND L3
     FILE 'CAPLUS' ENTERED AT 12:39:45 ON 29 JUN 2004
          47408 BROMINE
L5
          66442 ISOCYANATE
L6
            197 L5 AND L6
L7
             67 L5(L)L6
\Gamma8
          12353 PHOSGENE
L9
L10
             11 L7 AND L9
L11
           4350 L1
L12
           1655 L1/PREP
             18 L9 AND L12
L13
                SAVE TEMP ALL ISOCYANSRCH/L
=> color
        374523 COLOR
         39522 COLORS
        394562 COLOR
L14
                 (COLOR OR COLORS)
=> 17 and 1k14
             0 LK14
             0 L7 AND LK14
L15
=> 17 and 114
L16
             1 L7 AND L14
=> d 116
L16 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2004 ACS on STN
AΝ
     2000:455825 CAPLUS
     133:222175
DN
     High Loading Cellulose Based Poly(alkenyl) Resins for Resin Capture
ΤI
     Applications in Halogenation Reactions
     Chesney, Antony; Steel, Patrick G.; Stonehouse, David F.
ΑU
     Department of Chemistry, University of Durham Science Laboratories,
CS
     Durham, DH1 3LE, UK
SO
     Journal of Combinatorial Chemistry (2000), 2(5), 434-437
     CODEN: JCCHFF; ISSN: 1520-4766
PB
     American Chemical Society
DΤ
     Journal
LA
    English
RE.CNT 30
              THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS RECORD
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
=> pha
         10389 PHA
           974 PHAS
L17
         11033 PHA
                 (PHA OR PHAS)
```

=> apha

1160 APHA

4 APHAS

L18

1163 APHA

(APHA OR APHAS)

=> 17 and 118

L19

0 L7 AND L18

=> logoff hold

COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	61.87	80.08
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL

ENTRY SESSION
CA SUBSCRIBER PRICE -5.54 -5.54

SESSION WILL BE HELD FOR 60 MINUTES
STN INTERNATIONAL SESSION SUSPENDED AT 13:17:19 ON 29 JUN 2004

Connecting via Winsock to STN

Welcome to STN International! Enter x:x

LOGINID: SSSPTA1623PAZ

## PASSWORD:

\* \* \* \* \* RECONNECTED TO STN INTERNATIONAL \* \* \* \* \* \* SESSION RESUMED IN FILE 'CAPLUS' AT 13:38:01 ON 29 JUN 2004 FILE 'CAPLUS' ENTERED AT 13:38:01 ON 29 JUN 2004 COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	62.31	80.52
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	-5.54	-5.54
=> file reg		
COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	62.31	80.52
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	-5.54	-5.54

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Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 28 JUN 2004 HIGHEST RN 700803-86-7 DICTIONARY FILE UPDATES: 28 JUN 2004 HIGHEST RN 700803-86-7

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2004

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

```
=> e isophorone diamine/cn
                   ISOPHORONE DIACETOACETAMIDE/CN
E2
             1
                   ISOPHORONE DIACETOACETAMIDE-TRIPROPYLENE GLYCOL DIACRYLATE C
                   OPOLYMER/CN
E3
             1 --> ISOPHORONE DIAMINE/CN
E4
                   ISOPHORONE DIAMINE-1,1'-METHYLENEBIS(4-ISOCYANATOCYCLOHEXANE
                   )-POLYETHYLENE GLYCOL BLOCK COPOLYMER/CN
E5
             1
                   ISOPHORONE DIAMINE-2-ETHYLHEXYL GLYCIDYL ETHER COPOLYMER/CN
                   ISOPHORONE DIAMINE-IPDI-KURAPOL P 2010 BLOCK COPOLYMER/CN
E6
             1
E7
             1
                   ISOPHORONE DIAMINE-IPDI-POLYOXYPROPYLENE-PLACCEL CD 220PL CO
                   POLYMER/CN
             1
                   ISOPHORONE DIAMINE-ISOPHORONE DIISOCYANATE-2,2,4-TRIMETHYL-1
E.8
                   ,6-HEXANEDIOL-2,4,4-TRIMETHYL-1,6-HEXANEDIOL COPOLYMER/CN
             1
                   ISOPHORONE DIAMINE-ISOPHORONE DIISOCYANATE-JEFFAMINE ED 900
E9
                   BLOCK COPOLYMER/CN
E10
             1
                   ISOPHORONE DIAMINE-ISOPHORONE DIISOCYANATE-PHTHALIC ANHYDRID
                   E COPOLYMER/CN
E11
             1
                   ISOPHORONE DIAMINE-ISOPHORONE DIISOCYANATE-POLY(TETRAMETHYLE
                   NE GLYCOL)-TETRAMETHYLXYLYLENE DIISOCYANATE COPOLYMER/CN
                   ISOPHORONE DIAMINE-ISOPHORONE DIISOCYANATE-PROPOXYLATED BISP
E12
                   HENOL A-TEREPHTHALIC ACID-TRIMELLITIC ANHYDRIDE COPOLYMER/CN
=> e3
             1 "ISOPHORONE DIAMINE"/CN
L20
=> d 120
L20 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2004 ACS on STN
    2855-13-2 REGISTRY
RN
     Cyclohexanemethanamine, 5-amino-1,3,3-trimethyl- (9CI)
                                                            (CA INDEX NAME)
CN
OTHER CA INDEX NAMES:
     Cyclohexanemethylamine, 5-amino-1,3,3-trimethyl- (7CI, 8CI)
OTHER NAMES:
CN
     1,3,3-Trimethyl-1-aminomethyl-5-aminocyclohexane
CN
     1-Amino-3,3,5-trimethyl-5-aminomethylcyclohexane
CN
     1-Amino-3-(aminomethyl)-3,5,5-trimethylcyclohexane
CN
     3,3,5-Trimethyl-5-aminomethylcyclohexylamine
CN
     3-Aminomethyl-3,5,5-trimethylcyclohexylamine
CN
     5-Amino-1,3,3-trimethylcyclohexanemethanamine
     5-Amino-1,3,3-trimethylcyclohexanemethylamine
CN
CN
    Araldite HY 5083
CN
    Chemammina CA 17
CN
    Epilox H 10-31
CN
    IPD
CN
    IPDA
CN
    Isophorone diamine
CN
    Luxam IPD
CN
    Polypox IPD
CN
    Rutadur SG
```

CN Vestamin IPD

FS 3D CONCORD

DR 177646-11-6, 129050-51-7, 25495-81-2, 50858-71-4, 52004-55-4, 45981-71-3, 52697-24-2, 116723-72-9

MF C10 H22 N2

CI COM

LC STN Files: ANABSTR, AQUIRE, BEILSTEIN\*, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CAOLD, CAPLUS, CASREACT, CBNB, CHEMCATS, CHEMLIST, CHEMSAFE, CIN, CSCHEM, CSNB, EMBASE, HSDB\*, IFICDB, IFIPAT, IFIUDB, MEDLINE, MSDS-OHS, NIOSHTIC, PIRA, PROMT, RTECS\*, TOXCENTER, ULIDAT, USPAT2, USPATFULL (\*File contains numerically searchable property data)

Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

DT.CA Caplus document type: Conference; Journal; Patent; Report

RL.P Roles from patents: BIOL (Biological study); FORM (Formation, nonpreparative); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses); NORL (No role in record)

RLD.P Roles for non-specific derivatives from patents: BIOL (Biological study); MSC (Miscellaneous); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)

RLD.NP Roles for non-specific derivatives from non-patents: BIOL (Biological study); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)

## \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

1226 REFERENCES IN FILE CA (1907 TO DATE)
612 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
1226 REFERENCES IN FILE CAPLUS (1907 TO DATE)
8 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

=> file caplus COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 7.04 87.56 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE 0.00 -5.54

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FILE COVERS 1907 - 29 Jun 2004 VOL 141 ISS 1 FILE LAST UPDATED: 28 Jun 2004 (20040628/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> 120/prep

1226 L20

3164945 PREP/RL

L21

369 L20/PREP

(L20 (L) PREP/RL)

=> formaldehyde

130321 FORMALDEHYDE

368 FORMALDEHYDES

L22 130427 FORMALDEHYDE

(FORMALDEHYDE OR FORMALDEHYDES)

=> 121 and 122

L23 13 L21 AND L22

=> aniline

93295 ANILINE

12073 ANILINES

L24 98085 ANILINE

(ANILINE OR ANILINES)

=> 123 and 124

L25 3 L23 AND L24

=> d 125 1-3 ti fbib abs

L25 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2004 ACS on STN

TI Production of amine-formaldehyde condensation products

AN 2001:903322 CAPLUS

DN 136:38254

TI Production of amine-formaldehyde condensation products

IN Stroefer, Eckhard; Mueller, Christian; Sohn, Martin; Kaibel, Gerd

PA Basf A.-G., Germany

SO Ger. Offen., 8 pp. CODEN: GWXXBX

DT Patent

LA German

FAN.CNT 1

PΙ

PATENT NO. KIND DATE APPLICATION NO. DATE

DE 10027778 A1 20011213 DE 2000-10027778 20000607

DE 2000-10027778 20000607

AB Amine-formaldehyde condensation products are obtained by conversion of at least one amine (A) with a mixture (B) of

poly(oxymethylene) glycol, HCHO monomer, methylene glycol, and water, characterized in that a fractionation of mixture B and the conversion with amine A takes place in a reaction column, whereby amine A and the portion of the fractionated mixture B reacting with the amine A move countercurrently with each other. This method is especially useful in

producing
 methylenedianiline from PhNH2 as amine A with reduced N-methylated
 byproducts.

- L25 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Centipede polymers grafted with hydrogenated block copolymers and polyalkylenes and gels thereof
- AN 1999:722767 CAPLUS
- DN 131:337863
- TI Centipede polymers grafted with hydrogenated block copolymers and polyalkylenes and gels thereof
- IN Wang, Xiaorong; Matsuse, Takahiro; Foltz, Victor J.; Mashita, Naruhiko; Hall, James E.; Toyosawa, Shinichi; Takeichi, Hideo
- PA Bridgestone Corporation, Japan
- SO Eur. Pat. Appl., 16 pp. CODEN: EPXXDW
- DT Patent
- LA English
- FAN.CNT 1

FAM.CMI I			
PATENT NO.	KIND DATE	APPLICATION NO.	DATE
PI EP 955329	A1 19991110	EP 1999-107308	19990419
R: AT, BE,	CH, DE, DK, ES, FR,	GB, GR, IT, LI, LU,	, NL, SE, MC, PT,
IE, SI,	LT, LV, FI, RO		
		US 1998-73617 A	19980506
US 6054532	A 20000425	US 1998-73617	19980506
JP 11343320	A2 19991214	JP 1999-120365	19990427
		US 1998-73617 A	19980506
CA 2270372	AA 19991106	CA 1999-2270372	19990428
		US 1998-73617 A	19980506

- AB The present invention teaches a method for enabling the formation of a high damping, soft polymer gel. The method includes: reacting a alkenylbenzene-maleimide copolymer with a maleated polyalkylene and a maleated hydrogenated block copolymer and an alkyl diamine grafting agent under substantially dry conditions sufficient to form a hydrogenated block copolymer-polyalkylene grafted poly(alkenyl benzene-co-maleimide) polymer product, and dispersing this product with an extender oil sufficient to form the gel.
- RE.CNT 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L25 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Two-component castor oil- and polyoxyalkylene-polyurethane adhesives dispensable in 1:1 volume-ratio.
- AN 1995:990702 CAPLUS
- DN 124:30670
- TI Two-component castor oil- and polyoxyalkylene-polyurethane adhesives dispensable in 1:1 volume-ratio.
- IN Trinks, Rainer; Stepanski, Horst; Colinas-Martinez, Jose; Ganster, Otto
- PA Bayer A.-G., Germany
- SO Eur. Pat. Appl., 8 pp. CODEN: EPXXDW
- DT Patent
- LA German
- FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

```
PΙ
     EP 676427
                       A2
                            19951011
                                           EP 1995-104284
                                                            19950323
     EP 676427
                      A3
                            19960605
         R: DE, FR, GB, IT
                                           DE 1994-4411666 19940405
     DE 4411666
                            19951012
                                           DE 1994-4411666 19940405
                       A1
     CA 2146076
                       AA
                            19951006
                                           CA 1995-2146076 19950331
                                           DE 1994-4411666 19940405
     JP 07278518
                            19951024
                       A2
                                           JP 1995-97665
                                           DE 1994-4411666 19940405
    Polyurethane adhesives dispensable in a 1:1 volume ratio consist of: (1) a
     polyisocyanate with a NCO content of 10-25 weight%, a maximum viscosity of 2500
     mPa-s (at 25°), and 2-3 average functionality, (2) an aliphatic diol. with
     an OH number of >835 mg KOH/g, (3) a polyol with an average OH number of
20-200 mg
     KOH/g and 2-4 average functionality, and (4) at least one aliphatic or
aromatic di-
     or trifunctional amine, with maximum mol. weight 300 g/mol. The total average
OH
     number and maximum viscosity of components 2-4 are 200-290 mg KOH/g and 3000
     mPa-s (at 25°), resp., and the diol, polyol, and amine components
     do not sep. from the mixture The compns. can also contain 0.005-2.0 weight%,
     based on total adhesive weight, of a suitable tertiary amine catalyst. The
     polyisocyanate component (component 1) and the diol-polyol-amine component
     (components 2-4) are packaged sep. and then dosed in a 1:1 volume ratio from
     sep. containers.
=> d his
     (FILE 'HOME' ENTERED AT 12:35:23 ON 29 JUN 2004)
     FILE 'REGISTRY' ENTERED AT 12:35:51 ON 29 JUN 2004
               E ISOPHORONE DIISOCYANATE/CN
              1 E3
L1
     FILE 'REGISTRY' ENTERED AT 12:38:33 ON 29 JUN 2004
L2
          2357 BROMINE
L3
          17997 ISOCYANATE
L4
              7 L2 AND L3
     FILE 'CAPLUS' ENTERED AT 12:39:45 ON 29 JUN 2004
L5
          47408 BROMINE
          66442 ISOCYANATE
L6
L7
           197 L5 AND L6
L8
             67 L5(L)L6
L9
          12353 PHOSGENE
L10
            11 L7 AND L9
L11
           4350 L1
          1655 L1/PREP
L12
L13
             18 L9 AND L12
                SAVE TEMP ALL ISOCYANSRCH/L
L14
         394562 COLOR
              0 L7 AND LK14
L15
L16
              1 L7 AND L14
L17
          11033 PHA
          1163 APHA
L18
L19
             0 L7 AND L18
     FILE 'REGISTRY' ENTERED AT 13:38:12 ON 29 JUN 2004
               E ISOPHORONE DIAMINE/CN
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L20

1 E3

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FILE 'CAPLUS' ENTERED AT 13:39:11 ON 29 JUN 2004
L21
            369 L20/PREP
        130427 FORMALDEHYDE
L22
L23
            13 L21 AND L22
         98085 ANILINE
L24
L25
             3 L23 AND L24
=> 120/rct
         1226 L20
      2634689 RCT/RL
L26
          206 L20/RCT
                 (L20 (L) RCT/RL)
=> 113 and 126
          13 L13 AND L26
L27
=> 109 \text{ and } 127
             1 LO9
L28
             0 LO9 AND L27
=> 19 and 127
           13 L9 AND L27
L29
=> d 29 1-13 ti fbib abs
     13 ANSWERS ARE AVAILABLE. SPECIFIED ANSWER NUMBER EXCEEDS ANSWER SET SIZE
The answer numbers requested are not in the answer set.
ENTER ANSWER NUMBER OR RANGE (1):end
=> d 127 1-13 ti fbib abs
L27 ANSWER 1 OF 13 CAPLUS COPYRIGHT 2004 ACS on STN
    Improved process for the preparation of di- and triisocyanates by
     continuous phosgenation of amines
    2003:470304 CAPLUS
AN
DN
    139:36959
    Improved process for the preparation of di- and triisocyanates by
     continuous phosgenation of amines
    Friedrich, Martin; Stutz, Herbert
IN
PΑ
    Bayer AG, Germany
    Eur. Pat. Appl., 6 pp.
SO
    CODEN: EPXXDW
DT
    Patent
    German
LA
FAN.CNT 1
                 KIND DATE
                                          APPLICATION NO. DATE
    PATENT NO.
                           -----
                     ____
                                          _____
    EP 1319655
EP 1319655
                     A2
                           20030618
                                          EP 2002-26860 20021202
                    A3 20031210
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, SK
                                          DE 2001-10161384A 20011214
    DE 10161384
                      A1
                           20030618
                                          DE 2001-10161384 20011214
    US 2003114705
                      A1
                           20030619
                                          US 2002-316749 20021211
                                          DE 2001-10161384A 20011214
     JP 2003192658
                     A2
                           20030709
                                          JP 2002-360733 20021212
                                          DE 2001-10161384A 20011214
    CN 1425647
                      Α
                           20030625
                                          CN 2002-157003 20021216
                                          DE 2001-10161384A 20011214
    MARPAT 139:36959
```

OS

Di- and triisocyanates R(NCO)n [R = (cyclo)aliphatic or aromatic C≤15 hydrocarbon residue, with a proviso; n = 2, 3] are manufactured by continuous phosgenation of di- and triamines R(NH2)n (R, n as defined). The vapors

of di- or triamines, optionally diluted with inert gas or inert solvent vapors, are preheated to 200-600° and introduced into a static mixer in a tubular reactor, where they are mixed with preheated (200-600°) COCl2 which is introduced sep. The mixer has a specified geometry. Thus, isophorone diisocyanate was manufactured in 98.8% yield from 1:4:0.1 mol. mixture of isophoronediamine, COCl2 and N.

```
ANSWER 2 OF 13 CAPLUS COPYRIGHT 2004 ACS on STN
     Production of isocyanates in the gaseous phase
AN
     2003:434519 CAPLUS
DN
     139:22615
ΤI
     Production of isocyanates in the gaseous phase
     Woelfert, Andreas; Mueller, Christian; Stroefer, Eckhard; Pfeffinger,
IN
     Joachim; Weber, Markus; Knoesche, Carsten
     BASF Aktiengesellschaft, Germany
PA
     PCT Int. Appl., 21 pp.
SO
     CODEN: PIXXD2
DT
     Patent
LА
     German
FAN.CNT 1
     PATENT NO.
                     KIND DATE
                                          APPLICATION NO. DATE
     ___________
    WO 2003045900
PΙ
                     A1 20030605
                                         WO 2002-EP12930 20021119
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
             PL, PT, RO, RU, SC, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT,
             TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ,
            MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG,
             CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
             PT, SE, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR,
             NE, SN, TD, TG
                                           DE 2001-10158160A 20011128
    DE 10158160
                            20030612
                                          DE 2001-10158160 20011128
                      A1
     The invention relates to a method for producing diisocyanates by reacting
AB
    primary diamines with phosgene in the gaseous phase. Said
    method is characterized in that the reaction of diamine and
    phosgene occurs in a reaction channel, the internal dimensions of
    which have a width/height ratio of at least 2/1. With these dimensions
     the reaction chamber is useful for a longer period of time before it is
     necessary to clean the chamber of solid precipitate
              THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 2
             ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 3 OF 13 CAPLUS COPYRIGHT 2004 ACS on STN
TΙ
     Evaporative removal of aliphatic and cycloaliphatic isocyanate monomers
     from polymeric residues
ΑN
     1999:635503 CAPLUS
DN
     131:243741
     Evaporative removal of aliphatic and cycloaliphatic isocyanate monomers
ΤI
     from polymeric residues
    Mason, Robert W.; Fadakar, Farhad; Bridges, Joseph P.; Butler, Larry K.;
IN
     Keyvani, Majid
     Arco Chemical Technology, L.P., USA
PA
SO
     U.S., 9 pp.
     CODEN: USXXAM
DT
     Patent
```

FAN.CNT 1

English

LΑ

PATENT NO. KIND DATE

APPLICATION NO. DATE

PI US 5962728 A 19991005 US 1997-961800 19971031 US 1997-961800 19971031

AB A process for isolating aliphatic (e.g., 1,6-diisocyanatohexane) or cycloaliph. isocyanate monomer(s) from a liquid or viscous paste composition containing polymeric isocyanate residues and the isocyanate monomer(s) comprises: (A) introducing the composition into the heating zone of a dispersing evaporative dryer which contains both a heating and a cooling zone; (B) heating the composition to a temperature sufficient to cause monomer evaporation

forming a gaseous stream of isocyanate monomer(s), which is condensed and collected, and a molten stream of polymeric residue byproduct; and (C) moving the molten residue stream to the cooling zone in the dispersing evaporative dryer to cause solidification, forming a solid polymeric isocyanate residue having an isocyanate monomer content of <1%.

RE.CNT 18 THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L27 ANSWER 4 OF 13 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Purification of aliphatic isocyanates
- AN 1996:35267 CAPLUS
- DN 124:201653
- TI Purification of aliphatic isocyanates
- IN Nozawa, Kaneo; Matsuhira, Nobuya; Naito, Taketoshi; Morinaka, Katsutoshi; Tabuchi, Toshihiko
- PA Showa Denko Kk, Japan
- SO Jpn. Kokai Tokkyo Koho, 7 pp. CODEN: JKXXAF
- DT Patent
- LA Japanese

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	JP 07278088	A2	19951024	JP 1994-74915	19940413
	JP 2915784	B2	19990705		

AB The process comprises heating solns. of hydrolyzable Cl-containing crude aliphatic isocyanates and inert organic solvents at 140-270°, optionally mixing the crude isocyanates with the solvents at one time or gradually, distilling away higher amts. of the solvents than the weight of the isocyanates contained for ≥2 h, and optionally distilling the isocyanates from residue. A solution of isophorone diisocyanate (I) and 423 ppm hydrolyzable Cl in o-Cl2C6H4 was distilled at 175-185° and 560-600 mm Hg for 3 h to distill away o-Cl2C6H4, then distilled at 127° and 2 mm Hg to give 36 ppm hydrolyzable Cl-containing I with Harzen color number ≤10.

JP 1994-74915

19940413

- L27 ANSWER 5 OF 13 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of aliphatic polyisocyanates from polyamines and phosqene
- AN 1995:992543 CAPLUS
- DN 124:88109
- TI Preparation of aliphatic polyisocyanates from polyamines and
- PA Mitsui Toatsu Chemicals, Inc., Japan
- SO Ger. Offen., 11 pp.

CODEN: GWXXBX

- DT Patent
- LA German

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	DE 19510259	A1	19950928	DE 1995-19510259	19950321

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DE 19510259
                  C2
                       19970904
                                      JP 1994-50082 A 19940322
JP 07309827
                                      JP 1995-42956
                  A2
                       19951128
                                                     19950302
JP 3201921
                  B2
                       20010827
                                      JP 1994-50082 A 19940322
US 5523467
                       19960604
                                      US 1995-401807 19950310
                  Α
                                      JP 1994-50082 A 19940322
CN 1125718
                  Α
                       19960703
                                      CN 1995-104546
                                                     19950322
CN 1062857
                       20010307
                  В
                                      JP 1994-50082 A 19940322
In the conversion of an aliph polyamine to the polyisocyanate (e.g.,
m-xylylenediamine to m-xylylene diisocyanate) in an inert liquid medium, an
inert gas is added to the reactor during the reaction to increase the
yield of polyisocyanate and reduce the amount of phosgene
required.
ANSWER 6 OF 13 CAPLUS COPYRIGHT 2004 ACS on STN
Preparation of isophorone diisocyanate
1995:986708 CAPLUS
```

TI

ΑN

DN124:57000

TIPreparation of isophorone diisocyanate

IN Suguro, Yoshio; Katogi, Mamoru; Matsumoto, Masashi

Mitsubishi Kagaku KK, Japan PΑ

SO Jpn. Kokai Tokkyo Koho, 6 pp.

CODEN: JKXXAF

DTPatent

Japanese LА

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	JP 07252200	A2	19951003	JP 1994-44139	19940315
				JP 1994-44139	19940315

AB The title compound (I) is prepared from isophoronediamine (II) using continuous multireactors having ≥2 direct-binding phosgenating baths followed by a hydrochloride salt-forming bath, in which II is treated with HCl in inert solvents in the hydrochloride salt-forming bath under high temperature, the slurry obtained is phosgenated in the phosgenating baths, then the hydrochloride salt-forming bath is cooled to continue reaction after the phosgenation giving no ppts. A reactor having 4 baths, in which HCl was fed into 1st bath at 100° and COCl2 was fed into 2nd and 3th baths at 130° and 140°, resp., was fed with a solution of II in decalin into 1st bath to give I. The same treatment was carried out after no precipitate formation at 78° in the 1st reactor to give I without precipitation

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L27 ANSWER 7 OF 13 CAPLUS COPYRIGHT 2004 ACS on STN
```

TIPreparation of diisocyanates without using phosgene

AN 1994:701575 CAPLUS

DN 121:301575

Preparation of diisocyanates without using phosgene TI

Yanagii, Toyokazu; Itokazu, Teruo; Oka, Kenji IN

Daicel Chem, Japan PA

SO Jpn. Kokai Tokkyo Koho, 10 pp.

CODEN: JKXXAF

DTPatent

LΑ Japanese

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	JP 06172292	A2	19940621	JP 1993-218747 JP 1993-218747	19930902 19930902
				OF T333-7T0141	エフフンひりひと

Diisocyanates, useful for polyurethane manufacture (no data), are prepared by AΒ (a)

reaction of CO, O, and MeOH, (b) reaction of the resulting Me2CO3 with diamines in the presence of alkaline catalysts, and (c) pyrolysis of the resulting urethanes in the presence of catalysts at 1-700 Torr. Thus, N (sic), CO, and Ar/O were introduced to MeOH containing PdCl2, AcoCu and MgCl2 at 130° for 1 h and resulting Me2CO3 was treated with isophoronediamine and MeONa in MeOH at 70° for 6 h to give 99.5% isophorone dicarbamate, which was heated in dibenzyltoluene with Mn acetate under reflux at 10 Torr to give 74% isophorone diisocyanate.

- L27 ANSWER 8 OF 13 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of isophorone diisocyanate from isophoronediamine
- AN 1993:650204 CAPLUS
- DN 119:250204
- TI Preparation of isophorone diisocyanate from isophoronediamine
- IN Suguro, Yoshio; Kawamura, Shigenori
- PA Mitsubishi Chemical Industries Co., Ltd., Japan
- SO Jpn. Kokai Tokkyo Koho, 3 pp. CODEN: JKXXAF
- DT Patent
- LA Japanese
- FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	JP 05065265	A2	19930319	JP 1991-229135	19910909
				JP 1991-229135	19910909

- OS CASREACT 119:250204
- AB Isophorone diisocyanate (I) is prepared by reacting isophoronediamine or its hydrochloride salt (II) with **phosgene** in an inert organic solvent followed by distilling off the solvent and treating the crude product at 160-170° in an atmospheric of inert gas. Thus, II was treated with **phosgene** in decane and the reaction mixture was distilled to give crude I, which, after gel permeation chromatog., a product containing 4.2% impurities. This product was passed through a ball filter under the introduction of nitrogen gas at 1.9 L/h at 220° for 1 h to give a product containing 3.8% impurities.
- L27 ANSWER 9 OF 13 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Process for preparing polyurethanes for coatings
- AN 1992:61691 CAPLUS
- DN 116:61691
- TI Process for preparing polyurethanes for coatings
- IN Yagii, Toyokazu; Maruyama, Toshihide; Murata, Kiyokazu
- PA Daicel Chemical Industries, Ltd., Japan
- SO PCT Int. Appl., 77 pp.
- CODEN: PIXXD2
- DT Patent
- LA Japanese
- FAN.CNT 1

	PATENT NO.	KIND DATE	AP	PLICATION NO.	DATE
ΡI	WO 9114725 W: US	A1 1991	1003 WO	1991-JP369	19910319
	RW: CH, DE,	FR, GB, IT,	NL		
			JP	1990-68643 A	19900319
			JP	1990-76098 A	19900326
			JР	1990-88046 A	19900402
			JP	1991-99876 A	19910201
	JP 03275661	A2 1991	1206 JP	1990-76098	19900326
	JP 03287570	A2 1991	1218 JP	1990-88046	19900402
	JP 2997501	B2 2000	0111		
	JP 05262715	A2 1993	1012 JP	1991-99876	19910201
	JP 04211481	A2 1992	0803 JP	1991-52667	19910318

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JP 1990-68643 Al19900319
EP 477376
                 A1 19920401
                                     EP 1991-906281 19910319
    R: CH, DE, FR, GB, IT, LI, NL
                                     JP 1990-68643 A 19900319
                                     JP 1990-76098 A 19900326
                                     JP 1990-88046 A 19900402
                                     JP 1991-99876 A 19910201
                                    WO 1991-JP369 W 19910319
US 5138015
                 Α
                      19920811
                                    US 1991-752481 19910906
                                     JP 1990-68643 A 19900319
                                     JP 1990-76098 A 19900326
                                     JP 1990-88046 A 19900402
                                     JP 1991-99876 A 19910201
                                    WO 1991-JP369 W 19910319
The title process comprises preparing a dialkyl carbonate without using
```

The title process comprises preparing a dialkyl carbonate without using phosgene, reacting the carbonate with a diamine to give a urethane, thermally decomposing the urethane to form a diisocyanate, and reacting the diisocyanate with a polyol in the presence of a Lewis acid and/or a protonic acid to give a polyurethane. The polyurethane forms coatings with good heat and weather resistance. Isophorone diisocyanate (I) containing 0.1 ppm Cl was prepared by reacting CO in turn with MeOH and isophoronediamine (II), and decomposing the product. Heating I 44.4, polycaprolactone diol (PCl 220) 200, and dibutyltin dilaurate 0.046 g at 120° for 3 h, adding II 16.8, iso-Bu2NH 0.4, iso-BuCOMe 38.0, and iso-PrOH 199 g, and heating 3 h at 50° gave a solution having viscosity 304 P and containing 30.8% solids and <0.05% free NCO.

L27 ANSWER 10 OF 13 CAPLUS COPYRIGHT 2004 ACS on STN

TI Procedure for the production of (cyclo)aliphatic diisocyanates

AN 1989:231172 CAPLUS

DN 110:231172

TI Procedure for the production of (cyclo)aliphatic diisocyanates

IN Frosch, Hans Georg; Grave, Heinrich; Stutz, Herbert; Waldau, Eckart; Fuhrmann, Peter

PA Bayer A.-G., Fed. Rep. Ger.

SO Ger. Offen., 4 pp. CODEN: GWXXBX

DT Patent

LA German

FAN.CNT 1

	PATENT NO.	KIND DA	ATE AI	PPLICATION NO.	DATE
PI	DE 3714439 EP 289840 EP 289840 R: BE, DE,	A1 19 B1 19		2 1987-3714439 2 1988-106111	19870430 19880416
	22, 32,	25, 21,	•	1987-3714439	19870430
	US 4847408	A 19	9890711 US	1988-185721	19880425
	CA 1305165	A1 19		I 1987-3714439 A 1988-565025	19870430 19880425
	TD 62000050	20 40		1987-3714439	19870430
	JP 63280050 JP 08025984		9881117 JE 9960313	9 1988-104461	19880428
			DE	1987-3714439	19870430

OS CASREACT 110:231172; MARPAT 110:231172

AB A procedure for the preparation of OCNRNCO [R = C1-15(cyclo)aliphatic hydrocarbon

moiety] by phosgenation of the corresponding H2NRNH2 in the gas phase was characterized in that one: a) brings the gaseous diamine, optionally diluted with an inert gas or the vapors of an inert solvent, and COCl2, sep. heated to  $200\text{-}600^\circ$ , into reaction with each other in a cylindrical chamber at  $200\text{-}600^\circ$  without moving parts with the maintenance of a

turbulent streaming into the reactor chamber; b) leads the gas mixture which continuously leaves the reaction chamber through an inert solvent which is kept at a temperature above the decomposition temperature of the carbamoyl chloride

corresponding to the diamine; and c) subjects the diisocyanate dissolved in the inert solvent to a distillative work-up. In this manner, COC12 and H2N(CH2)6NH2 reacted at 400° to give 98.0% OCN(CH2)6NCO.

- L27 ANSWER 11 OF 13 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Multistep process for producing 3-isocyanatomethyl-3,5,5-trimethylcyclohexylisocyanate
- AN 1985:46407 CAPLUS
- DN 102:46407
- TI Multistep process for producing 3-isocyanatomethyl-3,5,5-trimethylcyclohexylisocyanate
- IN Hellbach, Hans; Merger, Franz; Towae, Friedrich
- PA BASF A.-G., Fed. Rep. Ger.
- SO Ger. Offen., 19 pp. CODEN: GWXXBX
- DT Patent
- LA German
- FAN.CNT 1

PA'	TENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI DE	3314790	A1	19841025	DE 1983-3314790	19830423
US	4596679	Α	19860624	US 1984-599821	19840413
				DE 1983-3314790	19830423
EP	126300	A1	19841128	EP 1984-104353	19840417
EP	126300	В1	19870401		
	R: BE, DE,	FR, GB	, IT, NL		
				DE 1983-3314790	19830423
CA	1225997	A1	19870825	CA 1984-452345	19840418
				DE 1983-3314790	19830423
JP	59205353	A2	19841120	JP 1984-78815	19840420
JP	05073737	B4	19931015		

DE 1983-3314790 19830423

- AB Isophorone diisocyanate (I) [4098-71-9] is prepared without the use of COCl2 by condensing isophoronediamine (II) [2855-13-2] with urea [57-13-6] and alcs. in the presence of carbonate and/or carbamate esters and, optionally, catalysts to give bis(alkoxycarbonyl) derivs. of II, separating and recycling the alcs. and esters, and cracking the II carbamate derivs. in the vapor phase. Thus, stirring II 1700, urea 1200, and BuOH [71-36-3] 300 g with (BuO)2CO [542-52-9] 105, H2NCO2Bu [592-35-8] 117, di-Bu isophoronedicarbamate (III) [78581-44-9] 956, and BuOH 3288 g (recovered from previous runs) at 210-220°/6-8 bar with NH3 distillation, stripping volatiles, volatilizing III at 270-280°/30 mbar, and cracking the vapors at 410° gave a mixture of I 78, monoisocyanate monocarbamates 19, and III 3%, distillation of which gave 1472 g I with purity >99%. The distilled volatiles and residues, containing BuOH 3151, (BuO)2CO
- H2NCO2Bu 113, and III 2066 g, were recycled.
- L27 ANSWER 12 OF 13 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Continuous preparation of 1-isocyanato-3-(isocyanatomethyl)-3,5,5-trimethylcyclohexane
- AN 1975:86817 CAPLUS
- DN 82:86817
- TI Continuous preparation of 1-isocyanato-3-(isocyanatomethyl)-3,5,5-trimethylcyclohexane
- IN Schmitt, Karl; Disteldorf, Josef; Reiffer, Johannes
- PA Veba-Chemie A.-G.
- SO Ger. Offen., 10 pp.

CODEN: GWXXBX

DTPatent LAGerman FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	DE 2323299	A1	19741121	DE 1973-2323299	19730509
	DE 2323299	C3	19831208		
	JP 50052048	A2	19750509	JP 1974-50377	19740508
	JP 58035179	B4	19830801		
				DE 1973-2323299	19730509
	US 3916006	Α	19751028	US 1974-467976	19740508
				DE 1973-2323299	19730509

1-Isocyanato-3-(isocyanatomethyl)-3,5,5-trimethylcyclohexane (I) AΒ [4098-71-9] was continuously prepared without agglomeration of the suspension by phosgenation of 1-amino-3-(aminomethyl)-3,5,5trimethylcyclohexane (II) [2855-13-2] in an inert solvent with excess COC12 at 130-60° in previously prepared I. Thus, COC12 [75-44-5] was passed into II in PhCl containing gaseous CO2 at 30°, the mixture passed into a reactor containing I in PhCl at 130° and then into a 2nd reactor for after reaction at 130° with passing of COC12 in countercurrent through both reactors to give 98% I of Cl content 0.1%.

L27 ANSWER 13 OF 13 CAPLUS COPYRIGHT 2004 ACS on STN

ΤI Isocyanates

ΑN 1973:431672 CAPLUS

DN 79:31672

TΙ Isocyanates

IN Edmondsen, John Neville; Hulse, Rae; Kerrigan, Vincent

PAImperial Chemical Industries Ltd.

SO Ger. Offen., 17 pp.

CODEN: GWXXBX

DTPatent

LΑ German FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO.	. DATE
PI DE 2249459 A1 19730419 DE 1972-2249459	20,21000
GB 1971-47795	19711014
IT 967968 A 19740311 IT 1972-29815	19720928
GB 1971-47795	19711014
NL 7213291 A 19730417 NL 1972-13291	19721002
GB 1971-47795	19711014
BE 789809 A1 19730406 BE 1972-122871	19721006
GB 1971-47795	19711014
FR 2157485 A5 19730601 FR 1972-36382	19721013
GB 1971-47795	19711014
JP 48048419 A2 19730709 JP 1972-103161	19721014
GB 1971-47795	19711014

AB Isocyanates were prepared by treating a mixture of (2) amines with COC12. Thus, a mixture of cyclohexylamine and a polyphenylamine, prepared by reacting HCHO and PhNH2, was treated with COCl2 to give cyclohexyl isocyanate and polyphenyl polyisocyanate; a mixture of n-C8H17NH2 and n-C18H37NH2 gave n-C8H17NCO and n-C18H37NCO.

A single name cannot be used for two saved items at the same time. Enter "Y" if you wish to replace the current saved name with a new definition. Enter "N" if the current saved definition must be preserved. You may then reenter the SAVE command with a different

<sup>=&</sup>gt; save temp all isocyansrch/l

<sup>&#</sup>x27;ISOCYANSRCH/L' IN USE

saved name. Enter "DISPLAY SAVED" at an arrow prompt (=>) to see a list of your currently defined saved names.

REPLACE OLD DEFINITION? Y/(N):y

L# LIST L1-L29 HAS BEEN SAVED AS 'ISOCYANSRCH/L'

=> logoff hold		
COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	61.13	148.69
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	-11.09	-16.63

SESSION WILL BE HELD FOR 60 MINUTES
STN INTERNATIONAL SESSION SUSPENDED AT 13:54:49 ON 29 JUN 2004